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- Solution phase nucleic acid sandwich assay and polynucleotids probes useful therein.
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0 225 80

Description

This invention relates generally to a solution phase nucleic acid sandwich assay.

Meinisch and Wah, Anz. Bischem., (1984) 1838/87-84, provide a moview article of hybridization for biotrispees. See also large yet at. Prov. BM. Acad. Sci. U.B. (1985) 2004-67-96, p. a description of the did biot saxys, Sandwich hybridization in described by Risele et al., Qur. Tep. Microbiol. Immunitory. (1985) pp. 506. The call the provided of the control of the

The increasing sear of cloning and synthesizing DNA sequences has greatly expanded opportunities for delacting particular nucleic seld sequences of Interest. No larger man one rely on the ord Immunocomplexes for the detection of pathogens, leutens, entigens, and the files. Rather than detecting particular determinant seas, one can offect DNA sequences or FRAA sequences associated with a particular cell. In this manner, diseases can be diagnosed, phenothypes and genotypes can be analyzed, as well as observablems. See the section of the control of the

For the most part, snelyses of DNA sequences have involved the binding of a sequence to e solid support and hybridization of a complementary sequence to the bound sequence. The enrositing and completing step usually knowns an astendic period of time and requires careful weaking to minimize nonse specific boolignound signals. There is substantial interest in developing new techniques for analyzing ruciols code sequences, which we more pedig, minimize the number of mediginally slages, and provide for on

Increased signed to notice ratio.

Operating European Patent Application No 60121564.8, which is divised from the present application is directed to polynomeocific probes useful in such inchrispes. The majority of polynomicotific probes in current one are reduced with belook of polynomicotific probes in current one are reduced with belook of polynomicotific probes with the contraction of th

radioactive are preferred.

so in order to incorporate other, non-rediscutive leges of detectable specieles in a ruciceoties, some sort of chemical modification of the medicated is required. It is which processive that ruciceoties modification is a difficult and sentitive procedure, as any modification reaction has to be mist enough to leven the RNA or DNA microscient states, while given a modified nucleoties product which can perfore in normal bases parting and stacking limerations. These consideration spicially finist mutucleos stabilishing positions to the value of the state of the

Oher considerations must also be base into account. Base painting may be liniteded during hybridization if the detectable label is a non-end of the nucleotide chain matter than present at some point with. Further, it has proved difficult to provide even non-radioschely labeled probes which may be inexpensively 4° synthesized in large outsethy. Thus, many known onches are limited in their obtainity associations.

Mentods and compositions are provided by the present invention to debucting particular multice and sequences. Two sets of magnetis are surpeplied, which are relation to as the capturing set and the labeling set. Each set has at least the members. The labeling set has (1) a first probe set, which completes one or a group of first analysis complementary asymment-first label renogration recognition respective consequence(s). The capturing or that of 10 a second probe to within completes one or a group of sequence(s) planed to associal capturing or than 6.11 a second probe to within completes one or a group of sequence(s) planed to a second capturing respect polytrucleotion recognition sequence(s). Given or a group of sequence(s) planed to a separate polytrucleotion recognition sequence(s) planed to a separate planed to the second capturing or the second capturing or the second capturing respect to the second capturing or the second capturing or the second capturing or the second capturing or the second capturing capturing capturing capturing capturing second capturing ca

The single stranded nucleic acid sample may be joined with the probes containing the complementary sequences of the two sets under annealing conditions, followed by the addition of the capturing and

optionally the labeling conjugates to provide for the analyte complex with the specific binding pair member and optionally the label. This probe hybridized analyte sequence is expansized by combrining the complex with the separating means and separating probe bound analyte from unbound analyte. When the label has not been previously added, the first recognition sequence-label conjugate is added to the phase containing it the securation or enable under the control in all the control i

The eforementioned copending European Patient Application No 90121594.8 describes modified derivatizable nucleotides having the structure of Formula 1, and their preparation including the step of derivations the RT molet with a detectable label:

Pormula 1

- wherein Pi is a reactive group derivatizable with a distinctive label, which reactive group ray to arrive, or cathogic of this did urther may be precised for virules upsticles managlations, Pi is an optional failing motivity such as those typically used to label proteins, and includes an amide, historite or distillated initiage or a combiation breach, Pi is selected from the gropp contesting of hydrogen, readly, thorrists, function and lodins, Pi is hydrogen, an anchoring group which covalently block the solutions to a solid support, or a blocking group and as dismolarly-pility or payl, which blocking group is gramming beneatable and advantage of the solid properties of the solutions to a solid support or a sense. Pi is hydrogen, an anchoring group which covalently block the solutions to a solid support or a sense. Pi is hydrogen, an anchoring group which covalently block the solutions to a solid support or a sense. Pi is a functional group usels are transfered explained in the proposition or the proposition of the proposit
- so The copending application also describes nucleic acid probes using one or more of the above modified nucleotides. The probe can be used to screen a sample containing a plurality of single-stranded or doublestranded polynucleotide chains and will label the desired sequence, if present, by hybridization.
 - In the accompanying drawing:-

10

Figure 1 is an illustrative depiction of a complex from the various components bound to a solid support (1) using DNA bridges for non-covalent binding and (2) using biotin-avidin bridges for non-covalent binding.

1. Sandwich Assay Method

Methods and compositions are provided for delecting a nucleic and sequence by employing two outs of requents. By unique contributions of models call designances complementary to a nucleic acid analyse and 5 to arbitrary sequences and specific binding pair members, a detectable bind may be sponsed into the phases in proportion to the amount of enable present in a sengle. By providing for amenting of uncleas and designance in solution, the time for performing the assay can be substantially diminished as compared to amenaling on a solid nuclear and the number of separations and weaking stays required can be limited to amenaling on a solid nuclear and the number of separations and weaking stays required can be limited to extend the second section of the design of the designation of the bindly instantiation of double animal dark of conductions are the section of the designation of the designation of conduction and the conduction of the designation of the bindly instantiation of double animal dark of conductions are the section of the section of the designation of the section of the designation of the designation of the properties of the section of the designation of the

concentrated form.

15 As indicated eleves, the method involves two sets of reagents. The first set results in labeling the analysis sequence. The second set provides the means for separating label bound to enalyte from unbound label in the assats medium.

The first set, the labeling set, will involve at least two reagents and may involve 10 to 00 resports or more. The first reagent will be a shorted nucleic acid regards and each enterher of the subset who so two nucleic soil regions. The first nucleic acid region of each member of the subset will be a region complementary to a sequence of the analysis. The second nucleotide sequence will be a recognision for the labeling regent. This second sequence will be selected, so as not to be encountered by endogenous sequences will be a tempolis.

The substits will have regions complementary to the satisfies sequence of at least 15 nucleotides (eff., 25 usually it less lettific, more usually a less less totte, and none than about 540, usually come the sequence of the sequence complementary to the shally the my be joined to a non-equentic sequence at either or both the 5 and 55-ments. The non-complementary sequence is judiciously relected no as not to brind to sequences in the assay which could result in fate positives, can be of any intent, usually fewer than middle.

o The complementary sequences will be chosen so as to leave areas for binding of the other reagents to the enalysis. Usually, areas of et least 25mt will be left available, where the analyse sequences complementary to the sequences of the individual members of the reagent subset may be substantially configuous or expansed and members of one subset may set when the sequence of the other subset. The particular pattern of binding between the two cubbets may vary which opposing on the sequences of the energy.

The reagent sequences may be prepared by synthesis in accordance with conventional procedures or by cloning and may be modified as appropriate for labelling.

The set of sequences which are complementary to the analyte may be selected based on a variety of consideration. Depending upon the native of the analyte, one may be interested in a consensus sequence, a sequence associated with polymorphisms, a particular phenotype or genotype, a particular strain, or the sell like. Thus, the labelling complementary sequences will be chosen in conjunction with the other complementary sequences of the conduring set to convoke information concerning the snath/hr.

The labeled sequence will include a expansor complementary to the first exception assumes of the labeling product, in labeling product, and included a labeling product and an include or seminated the labeling partially of labels. Windows expansion or may be present as a semination entertior or seminated the labeling partially of labels. Windows expansion of labeling products and labeling partially of labels. Windows labeling products and seminated labeling partial products and company fixed, Product has all company fixed, Product has all company fixed, Products and Company fix

Meinkoth and Wahl, And Bischem (1994) 132:287. The labels may be bound either covelently or noncovalently to the complementary sequence. Labels which may be employed include nationuclides, fluorescers, chemiluminescers, dyes, enzyme, enzyme substates, enzyme cotactors, enzyme shibitors, enzyme substates, enzyme corporation, and the like. Illustrative popolic labels include likeosecois, indomine. Treas red, privocyethid, unabellifices, luminol,

NADPH, o-f-galactocidate, horseradish peroxidate, etc.

The labeled sequence can be conveniently prepared by systitests. By providing for a terminal group which has a convenient functionality, various bobte may be joined through the functionality. Thus, one can provide for a carboxy, thiol, amine, hydrazine or other functionality to which the various lebels may be innered without detrimentally affecting duples formation with the sequence, has already included, one can

have a molecule with a plurality of labels joined to the sequence complementary to the labeling sequence. Alternatively, one may have a ligand bound to the labeling sequence and use a labeled receptor for binding to the linand to crowlet the labeling another complex.

This second set of reagenets provides the misens for separation of basis bound to analytis from vinbound to short. The misens for the separation or capitating mass inventives at least one capitating probe, usually a pluritify of probes defining a subset, which includes two polymochoolis sequence segions shill include a sequence of the labeling probe and a neopoliton sequence, offerent from the first about a recognition sequence of the labeling probe and separation of the labeling probe and separation above. The capitare probes may be between 10 the first set of recognition insists for the solicity growth and the sequence of the labeling probes the second advo. The capitaring sequences will be selected and synthesized in the same nancer as described above. The capitaring sequences will be selected and synthesized in the same capital problem. They have seme consistent with the labeling problem. They have some consistent with the labeling problem than of the same capital second with the labeling problem. They have seme consistent with the labeling problem, they have seme consistent with the inventor in propriating to approach to capitaring the seme consistent with the inventor in propriating the capitaring the second to the second section for the second section of the second section for the second section for the second section of the second section of the second section for the sec

While the separating means may be directly bound to a sequence complementary to the capturing in recognision sequence, perimetally a separation braining pair member will be bound to the complementary sequence. The appoint braining pair member will be a liquid or recognist, presented by aligned. Liquid smay be any miscrision for which a materially counting receptive effects or can be represent. These, materially or any miscrision of the sequence of the sequen

usually less than about 2,000 molecular weight, and preferably less than about 1,000 molecular weight.

The mospitors will generally be probin molecules and may include artibodies, naturally occurring proteins, such as avidin, thyroxine binding globular, etc., lectiles, enzymes, and the like. The receptors will generally be at least about 10,000 molecular weight, more usually 12,000 or more molecular weight, usually sets than about one million molecular weight.

The sportlic binding path member may be joined to the second recognition sequence by any convenient means. As sheaty indicated, the stoupones may be symbiotized, providing for a convenient structionality at the terminal base, which may then be used as the inkings bein shortly upon the particular binding pair members may be joined to the complementary sequence, despiriting upon the particular so choice or the specific binding pair members, its sites, and the nature of the functionalities. Althoratively, for a large specific binding pair members, the state of the functional time, and the nature of the functionalities. Althoratively, for a large specific binding pair members, the state of the specific binding pair members and the state of the functional state of the specific binding pair members are stated of the specific binding or conjugate with the proposed, so that them will be little interference, if any, from the specific binding with the little state of the displacement of the specific binding of the specific binding of the specific binding binding the binding binding the binding binding the binding pair members are specific binding and the specific binding binding the binding pair members are specific binding binding the binding pair members are specific binding binding the binding pair members are specific binding and the specific binding binding the binding pair members are specific binding binding the binding pair members are specific binding binding to be specific binding binding to be specific binding binding to be specific binding binding binding the binding binding binding pair members are specific binding binding binding binding pair members are specific binding bindin

Alternatively, the receptor may be an additional nucleotide sequence that specifically recognizes the recognition sequence of the capturing probe.

The separation means can be any support which allows for a rapid and clean separation of label bound to analyse from unboard ablot. Thus, the separation means may be particles, a solid wall surface of any of a variety of containers, e.g., certificing labeles, columns, immortate plate week, filters, buble, e.g.-Perdorably, op particles will be employed of a size in the range of about 0.4 to 200s, more usually from about 0.8 to 4.0s. The particles will be employed.

The homologous nucleic acid sequences need not have perfect complementarity to provide homoduplexes. In many situations, heteroduplexes will suffice where fewer then 15%, usually fewer than 10% of the bases are mismatches, incoming loops of five or more members.

48 Samples of assilyte nucleic acids may be from a variety of sources, e.g., biological fishiof or solids, do stiffs, environment materials, e.g., and may be proposed for the hybridization assilysis by a variety of means, e.g., probinsos KOSD, chardropic salls, etc. Alos, it may be of advantage to decrease the enverges, sociation, orbinical degradation (e.g., metal lond, etc. The fragments may be as small as O libb, usually belon at least final OSBs and may be at the or below.

in carrying out the method, the enables exequence will be provided in single standard form. Where the exequence is naturally present in estigle standard form, demanders will not be engined. However, where he exequence is naturally present in estigle standard form, demanders will not form the contract the execution of the provided standard form of the contraction of the contraction of the presence of the labeling probe entire the explaining probe, so that spon change of conditions to averailing conditions, the probles will standard contractions and contractions and contractions are contracting and contracting and contracting contracting and contracting contra

In many distallions, it will be preferable to avoid having either the bable or the separation manura present during destination. The elevated interpensature, the non-exposure solvents, the salts, or other materials present claring densitiarities may result in degradation, or undersinate modification of the label and/or separation manura. Therefore, in many silantions, densitiations may occur in the presence of the product separation coulding red ensembling of the product to the single-elevated OVM may occur. (Showed by the independent coulding radio elevation of the solutions are despressions, other makes or confirms, out as

Normally, the ratio of probe to anticipated moles of analytic will be at least 1:1, preferably at least about 1.5:1, and more preferably 2:1 and may be as high as 100:1 or higher. Concentrations of each of the probes 10 will generally range from about 10° to 10° ful, with sample nucleic acid concentrations varying from 10° 71.

After annealing conditions have been achieved, or even prior to such time, the labeled first recognition sequence and the capturing second encognition sequence are added and allowed to hybridize. Alternatively, the labeled first recognition sequence can be added after capture and separation.

A preferred embodiment which greatly reduces background and provides for extraordinarily high sonsitivity will employ the following sequence. With double-transfed sastyle, the enables will be denetured in the presence of the probe or complementary sequences, or the probe may be deded shortly other destauration, and under annositing conditions. After sufficient time for annosting, the complexes may the be combined with the secondaries means, whereby the considerate will be bound to the surrord. Any

consultation, and under almosting contained. After submoint time for amenting, the complexes may then be combined with the superation means, whereby the complexes will be bound to the support. Any so background DNA or non-specifically bound DNA may be washed sway so as to avoid non-specific binding of label in the next step. The solid support may then be washed to remove any non-specifically bound label to provide for a substentiely reduced background of non-specifically bound label.

Consider Figure 1, por 2. In effect, the another which is the long for at the top is combined with the A or 48 pictors, where A provider the complementary sequence for the label collegate and 8 provides are complementary sequence for the label collegate and 8 provides are complementary sequence for the specific brinding pair member, in this case, both. Thus, the A and 8 protein and the analysis would be joined dopather under sense(an) coulding, wherey complex formed would be considered under sense and provides and the analysis would be joined with the protein or the darked in a separate step to the solution containing the enalysis complement. After entitions they

for 6' to sensel to 8, the resulting biolinylated analyte complex would then be edided to the noticl support to which adds in 5 bound. After anticlinet time for the specific binding pair members to ferm complexes, the sold support could be washed free of any non-specific DNA, followed by the addition of the labeled sequence, which in this case is included as being fluorescent bound to A'. The bisheld sequence which in this case is included as being fluorescent bound to A'. The bisheld sequence being the specific control of the specific s

xx A comewhat shorter protocol is provided by the configuration depicted in part 1 of Figure 1. In this shatch, the protocal A and B would be added to the articles under amending conficion, whereby enterly complisses would form. After auditiont films for analytic complemes to form, the enterly complex sould now look of the based on the cold support for multical test for the opacting protocols to bird the section support by complex (immains with the sequence) self-cented and 5°C. Excess DNA could be weethed case, support to complex (immains with the sequence) self-cented and 5°C. Excess DNA could be writted case, support to complex (immains with the sequence) self-cented and 5°C. Excess DNA could be writted case, support to complex (immains to could be self-cented and the order. Excess in one-sectionity.)

bound label could then be washed away to provide the configuration depicted in Figure 1, part 1.

Usually, the dentaturing step will label from shout 5 to 25 minutes, usually from ebout 5 to 15 minutes, while the annealing step will generally take from about 30 minutes to 2 hours, frequency being completed in about 1 hour. Annealing can be carried out at a malfly elevated interperature, generally in the range from

about 20°C to 50°C, more usually srom ebout 25°C to 40°C, particularly 37°C.

Unsulty, on a squore medium is employed, particularly a latered superous medium, which may include various additives. Additives which may be employed include low concentrations of detergent (0,1 to 1%, saltx, e.g., socilum citats (0,017° to 0,170M). Root, polytripyproticions, carrier rusolici coids, carrier proteins, etc. Depending upon the nature of the specific briding pair members, various solvents may be

added to the aqueous medium, such as dimethytisomanide, dimethytisulfoode, and formanide. These other solvents will be present in amounts ranging from 2 to 50%. The stringency of the amessing medium may be controlled by temperature, sail concentration, solvent restren, and the like. Thus, depending upon the length and nature of the source of interest, the

For the separation step, for example, using a ligand-receptor pair, the medium may be changed to optimize or approximately optimize the conditions for specific binding pair complex formation. Thus, the pH will usually be modified to be in the range of about 6 to 9, preferably about 7. This can be readily schlewed.

by adding from about 0.5 to 2, usually about 1 volume of about a 0.1 to 0.5M buffered medium, e.g., properties buffered stallne, bit has annealing medium. This medium may be added in conjunction with expansion means and the mixture allowed to incubate for at least 5min, usually about 10min, and less than about 60min, usually about 15 to 45min, more usually about 00min being satisfactors.

5 The phases may then be expensed in accordance with the nature of the separation means. For particles, certification or Effective of provide for respectation of the particles, locating the supermatent or isolating the supermatent. Where the particles are assayed, the particles will be washed froughly, usually own one to five times, with an appropriate buffered medium, e.g., PSS. When the expension means the value of copport, the supermatent may be isolated or discarded and the wall washed in the same manner as it indicated for the acertises.

Depending upon the nature of the label, various techniques can be employed for detecting the presence of the label. For fluorescens, a large number of different fluoremeters are available. With enzymes, either e fluoremeter or a colored product can be provided and determined fluoremetrally, spectrophoto metrically or visually. The various labels which have been employed in lamsunossasy or after techniques 5 applicable to immunossasy can be employed with the soliced assers.

2. Nucleic Acid Probes

Nucleic edid probes useful in conjunction with the above assay method are probes which ere prepared from one or more modified nucleotides. As noted above, these nucleotides and probes are fully described in cooperfine European Patent Application No 80121594.8.

It is to be understood that while the invention has been described in conjunction with the preferred pacific embodiments thereof, the toroid project operations are sell as the Examples with follow or en intended to lituarists and not limit the scope of the invention. The following experimental and exemplifying materials includes, limit and and for convenience and completeness, description of nucleotides and probe perspection eithorized these appects even not aspects of the present invention as such. Again, the needer is referred to covereding Exception Pattern Application, the 50/2158445.

Experimental

Analyte	Bg.	lII	HBV	Frag	ment
---------	-----	-----	-----	------	------

ATCTCC	TAGACACCGCCTO ATCTGTGGCGGA	CAGCTCTG	TATCGAGAAG	CCT	TAGAGTCTCC	TGAC
AUU	NICIGIOGCOCN	I	AIMGUICIIC	J GGA	AICI CAGAGG	ACTO

CATTGCTCACCTCACC ATA CTGCACTCAGGCAAGCCATTCTCTCCTGCG GGG AATTGATG
GTAACGAGTGGAGTGG TAT GACGTGAGTCCGTTCGCTAAGAGACGACCC CCC TTAACTAC

ACTICTAGCTAGCTAGGTGGGTA ATA ATTIGGAAGATCCAGCATCTAGGGATCTIG TAG TAGTGAGATCGATGGACCCACCCAT TAT TAAACCTTCTAGGTCGTAGATCCCTAGAAC ATC AT

AATTATGTTAATACTAACGTGGGTTTAA AGA TCAGGCAACTATTGTGGTTTCATATATCT TTAATACAATTATGATTGCAGCCAAATT TCT AGTCGGTTGATAACACCAAAGTATATACA

T GCC TTACTTTTGGAAGAGAGACTGTACTTGAAT ATT TGGTCTCTTTTCCGAGTGTGCATT

AGACCCA TCTGCGTCTAG

+ indicates probed segments

5

50

1)

Labe.	lling and	Capturi	ing Probe S	ets (Refer	to Fig.
3*	GACTTOGA	CTTGGTCAA	ATCTCTCCC	GGAGTOGAGACATA:	ectette 5'
	CTCCTCAA	CACCTITCE B	TC ATCTCAGAG	GACTOCTAACGACTO 2	CACTOC!
	GACTTGCAJ	GTTGGTCAA	CT GACCTGACT	COUTTOGGTAAGAGA 3	CCACCC
	GTCGTCAAA	GACCTITCT B	IC TTANCTACT	GAGATOGATGGACO	ACCCAT
	GACTTGCAA	CTTCCTCAA	CT TAMACCTIC	TAGGTOGTAGATOCO 5	TAGAAC
	CTCCTCAAA	GAGGITTCT B	TC ATTTAATAC	AATTATGATTGCACG	CAAATT
	GACTTGCAA	GTIGGTCAA	ET ACTCCCTTC	ATAACACCAAAGTAT 7	ATAGAA
	CTGCTCAAA	CAGGITTCT B	C ATCANAC	ETTCTCTCTGACATG	AACTTA
	CACTTOCAA	PTTGGTCAA	+ ACCAGAGAA	AGOCTCACACCTAAG 9	CCTGAG
	GTCCTCAAA	CAGGITTCI	C CTCCCATATO	TOCTOCTTTACGGG	CATAGA
	GACTTGCAA	TTOCTCAM	T CTTCTCAAC	CCTTTGATGACAAC	AATCTG
	CTCCTCAAA	AGCTTTCTT	C COCTOCCTO	OCTOCAGGGGATCTT	CTTCTT
	+ Labeler recognition	Probe -	+ + Probe	Segments to HB	v +
	A - Fluor B - Biot	escein la	bel conjugat	e binding site nding site	

Label conliquite (A) for DNA or swider support:
Fiversection - 5 CTEAACGETCHACACATTCA.5'
46. DNA exquence (B'C) bound to solid support
5' GAAGABACCCTTTCACACACATGETCACAAAGGTTAACCATGTTTCTTGT.5'
Blofin conliquite (B) for swider support
Blofin - 5 CACCACTTTCTCCAMAGABAC 3'
Blofin - 5 CACCACTTCTCCCAMAGABAC 3'
Blofin - 5 CACCACTTCTCCCAMAGABC 3'
Blofin - 5 CACCACTTCTCCAMAGABC 3'
Blofin - 5 CACCACTTCCAMAGABC 3'
Blofin - 5 CACCACTTCCAM

Preparation of biotin or fluorescein labeled DNA (A' or B'):

N⁴ - (2" - aminoethyl) - decxycytosine - DNA

The salyfe is an IBV Bgll fragment as indicated above. (Valencelle et el. (1891) is Animal Virus 26 centels, et et Folde, B, Jandenk, P., Fou, C.F., Ausdanie Press, ins. N.Y. p. 59-710) is faithed in Initially and capturing probes are indicated, where 12 different sequences complementary to different sequences present in IBV are provided. Sor of the IBV complementary sequences in priced to a common sequence (A) for completing with the label conjugate (A). The other six IBV complementary sequences are joined to a common sequence (56 or completing with a biolitylatist sequence (57 or a third DM), as sequence (6°C) for Indring to a support. In Figure 1 is shown an Bustration of the final complete involving the IBV strate of the various resignation.

Example 1

Labeling of Caproic Acid Derivative

To 1 mmole of fluorescenie isothicognate in 5 ml of DMF was added 2 mmole of 8-minicognote and 54 ul of 94thylmines. After 54 h at from Interpretation, the product are soluted by proparation for larger chromatography (Winner and Logg, Isong, Clam. 181509 (1979)). The died product was supended; in 10 ml of 11 LIDMFTHE (AVI) to black 1.5 manuface IMF Mingroup seculational and i manuface of supended; not not in 10 ml of 11 LIDMFTHE (AVI) to black 1.5 member 6 Mingroup seculational and i manuface of supended; not not in 10 ml of 11 LIDMFTHE (AVI) to Mingroup seculation and 10 ml of 10

to

6-N*-(2-Aminoethyli- Deoxycytidin

Caruthors, supra; McBride and Caruthers, Tetrahedron Lett. 24:245 (1983)).

28 An alsylated derivative of descryotides, N°-Q-aminosthyl descryotidine (B) was prepared from properly protected descryotides view the electropid derivative as described by Presect on Ubsease, Testabeleon Lett. 21:2205 (1989). This latest derivative was convented in By of deplacement of the latest protective described view. 21:0205 (1999) and lategory controlly of the "Internation Lett. 22:3105 (1995). In latest consequenting of C-British Collegion and Letter of the Co

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Exemple 3

Probe Preparation (Fluorescein Label)

Synthetic oligonucleotides were prepared by an automated phosphoramidite method as described in Warner et et., DNA 3-401 (1984). Punification was carried out eccording to Sanchez-Pescedor and Urdee, DNA 3-399 (1984).

The eminorahyl definition of decopyolidine as programed in Example 2 was incorporated by standard coupling procedure during the eligentaceledia symbiletis and the puriside modified injectived design even used for incorporation of a florescenin blob is stolens. To a doed sample (A5 OO 200 units) of the aminorahyl decopyolidine containing deligenous views added 50 bit of DMF and 56 bit of 16 bit bit of

Example 4

Probe Preparation (Biotin Label)

Using the probes containing aminoethylcytidine as prepared in the previous example, biddin labeling as achieved as bidees. The oligonacideoide (6-5 CO 280 units) was taken up in 50 uii DMI soldium so phosphale, pri 70 and 50 uii of DMF soldien on oli of DMF soldien containing in grid or "long chain" N-ydroxysuccininnicity bident (Pierce Chamical) was added. After 18 h at room temperature, the bidinylated probe was purified as described for the floroscenia bidend probe.

Example 5

Preparation of Solid-Supported DNA Probe

5 Fingment SC is symthetic S0mely was 5-phosphoptated with 14-physicacided bissues and ATP using standard conditions. After galaxysisched was 6-beds per securious. Hydrocylistic lates (10mg, 0.81; Paredox Laboratorias) was washed with DMSO, then three protines of with MMSI (included of 5-phosphops) and the securious of 5-phosphops (10mg) and 10mg of 10m

Example 8

20 Assay for HBV DNA Using DNA Solid Support

A pBR322 clore containing the settler HBV genome (Velenzuels et al., Animal Vinus Genetics, R. Janichich, B. Riska and C.F. Froz, Edit, Academic-Pierro, Her Volgo, pp. 57-70 (1980) was cut with BgIII and used as the analyte nucleic soid. Analyte in 101m of formanide containing 6 piconoles or file telebeling and capturing probe set was beteated to 96° C for tribin and cooled to room temperature. To this inchinan, 62ul of water, 20ul of 20xSSC, 10ml of 1% NP40 and 2ul (10ug) of polyA are edited, vortexed and incubated at 37° C for 1ml.

The exist apported DNA probes (i) picomises 400gs (i) added and incubated for an additional 1.5th. The mithrate is contributed at 20,000g for 2-bins and the suppersuited discussed for the apport is varied for the property and the property of the property and the property of the propert

TABLE 1

		Fluorescence Counts	
45	Condition	(Average of 4)	
	0.5 pmole HBV	5062 +/- 345	
	0.25 pmole HBV	4117 +/- 262	
50	No Analyte	3197 +/- 520	
	No Biotinylated Probe	3856 +/- 642	

Assay for HBV DNA Using Avidin Support

5 Experiment 7a:

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Analytic was mixed and incubated with the labeling and capitaling probles as above. Birtin tabled proble (12 piccmoles) in Sail 14;0 was then added, vortexed and incubated at 37°C for 30min. To the militure, 20mil of a 0.25% (wky) 0.3a unitin later (Plandex Laborationes) in 15°CS is added and incubated at 37°C for 10 fb. The militure is washed, incubated with fluorescein probe, washed and read on the Plandex screen machine as deepshed above.

TABLE 2

	Fluorescence Counts
Condition	(Average of 4)
0.5 picomole HBV	4052 +/- 462
0.25 picomole HBV	2644 +/- 397
0.10 picomole HBV	1956 +/- 173
No Analyte	1641 +/- 370
No Biotinylated Probe	1631 +/- 474

Experiment 7b:

This HBV plasmid was conclusted to an average size of 500pp. The densturation and hybridization were carried out as above except that 30 picomoles of labeling and capturing probes were used and a 5h 40 censuling was employed. After incubstion with 30 picomoles of biblinylated probe (2h), 50,11 of 10,25% aridin bases were added and incubsted (1,5h). A Biodescrip probe was added and incubstion was carried out for 1h followed by washing and reading on the Soreen Machine so described above.

TABLE 3

		Fluorescence Counts
,	Condition	(Average of 4)
	0.5 picomole HBV	5748 +/- 244
	0.4 picomole HBV	5352 +/- 331
	0.3 picomole HBV	4716 +/- 243
	0.2 picomole HBV	4071 +/- 243
	0.1 picomole HBV	3320 +/- 271
,	No Analyte	1679 +/- 167
	No Biotinvlated Probe	1716 +/- 177

It is evident from the above results that a highly specific sensitive assay for specific muscles cod sequence is provided. Respect can be readly prepared to be expected or threshold. The related is a sequence in a sample determined. The method is entirely related to the sequence of the sensitive about the presents or diseased or asspectated in a sample determined. The method is provided to the sequence of the s

as Exemple 8

5'-Dimethoxytrityl-2'-Deoxyuridine

5 To 2-Decoyuridine (10 g. 44 minols) dried by coverposition of pyristine and suspended in pyristine (100 m) was added 18.4 g 64 minols) 4.4-dismborophythy chloride (IMIT-Q). The reaction was allowed to procosed for 18 h at room temperature, and 100 ml methanic was added to described excess DMIT-CI. Most of the pyristine was then removed in vacuou, and the residue, discovered in 500 ml eithy decisitie, was washed

with saturated aqueous NaHCO $_0$ (3x500 ml). The organic phase was dried over solid Na $_2$ SO $_4$ and eveporated to dryness. The residue was purified by flash chromatography on silica gel to give 18.0 g (77%) of 5'-dimethorythy's'-decorporations(C).

5 Example 9

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5'-O-(4,4'-Dimethoxytrityt)-3'-t-Butyldimethylsityt-2'-Deoxyuridine

To 18 g D4 mmoley of D is 200 nd DMF was added instances (S.B. g. S5 mmoley with guide stirling to sausure complete dissortion. Elluptimethylelly choices (S.B. g. 11 mmole) addressed in a small volume of 20 DMF was added doprelies with distring and the restriction was allowed to proceed in the disk for 18 in stroom temporation. The restriction nations was disduct with only access (200 mill good elected with NeWoCh. The restriction nation was discussed with a very considerable (200 mill good elected with NeWoCh. The restriction of the

(E)

4-(1,2,3,4-Tetrazoi-1-yi)-(5'-(4,4'-Dimethoxytrityi)-3'-t-Butyldimethytsityi-\$-D-2'-Deoxyribosyi] Pyridine-2(1H)-one

To 15.0 g (3 mmois) of D, dried by consposation of prividee and dissolved in pyridine (60 ml) was added diplentphosphate (2.6 g, 1.15 mmois) dissolved in pyridine (6 ml, 1-14 Meetitylene-2-europhyritatrization (MS-thyr) (15.5 g, 15.5 mmois) dissolved in pyridine (45 ml) was added and the rection mitures was already 25 ml writer. After 30 min, the product was concentrated under reduced pressure. The reducts was dissolved in 250 ml mellyhore choicked, washed with an exposure MS-FOD, soldine (CS25 ml), dried over dissolved in 250 ml mellyhore choicked, washed with an exposure MS-FOD, soldine (CS25 ml), dried over purpose the MS-FOD soldine (CS25 ml), dried the MS-FOD soldine (CS25

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4-N-(2-Amincethyl)-5'-Dimethoxytrityl-3'-t-Butyldimethylsityl-2'-Deoxycytidine

To a solution of athylene disentine (3.8 m.), 4.8 mmode) in disense (100 mil) cooled to 5.7 was added E (0.0 g. 4.3 mmole) and left for one bour. The solvent was removed at refeach greasure and the residue was conseporated with bulume to remove encous ethylene disense. The product was purified by — 5 chromotography on a silica gal couleme, nucled with 1.20 per method in mertiple coloristic by (girl 7.5 g. (76%)) of 4-Hr2-eminocophy6-5-dimentioncy(hy6-5-dimention(hy6-b)2-dimen

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(H)

N*-(N-FMOC-6-Aminocaproyl-2-Aminoethyll)-5'-Dimethyltrityl-3'-t-Butyldimethylsilyl-2'-Deoxycytidine

(CH,),-NH-FMOC

45 To a sainton of E (6.5 p. 8.8 monis) in pytidine (20 m) was added N+NOC-6-uninocaperia acid (4.26 p. 12 mmole) (PMOC represented by a motton 4) pas of DC (4.86 p. 4.8 mmole). Not 3 h, the necessities as judged by the (sites in 10% methanolimethylene chickels). Pytidine was removed at rectioned presents. The resistance sententiated with drifty acidine, incelluble dispolarization (ECHI) titlened of an of the solvent removed. The product was included by sites gad demandagority valued with 6% of the product of the product was included by the part demandagority valued with 6% of the product of the product was included by the part of the part of

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A solution of intributy/immonium hacide (15 mode, 15 ml of a 11% solution in TEP) and supusus IFE is (1.5 ml of a 50% separces solution) were included and bey consequention of pridrice. The residence idiaselved to profition (15 ml) and extend to (0.7 to 2.7 at mode) which was disasted by solutions. After 16 hours at 4°C the meation miduse was olitized with 20 ml mertylene cholicist. Concentration spusses, NAMOO, was cerefully decide followed by solid INMOOs, added gradually so as to meatitate the HPHyydrice. After directly one Very Solid. The organic places were acconstrated to an oil, with was subjected as to allow plot drawing the concentration of solid visits. All the concentration spusses, The product the VERMOO-E-minor corpors/seminoretyin/s-directly concentration spusses.

To 5.1 g (5.7 mmole) of I in methylene chloride containing (disopropylethylamine) was added

(chioro-N.N-discopony)samhomethosy phosphine, 1,3 mt [1.2 eq.], b) at 0°C under argon. After 1 hr, orbity acoste (300 mt) was added and washed with 80% saturated equence sodium chioride, effect right the organic phase over Ns,50s, the product in methylene chloride was added dropwise to hexane at -40°C to precipitate 4-3a (75%) of 1.4.

Example 15

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Synthesis of Horseradish Peroxidase (HRP): DNA Conjugates

Sequence 1 6°-LECATORACGITCHACCATTCHAC) where LAC = 10° (Braincoapprojulaminostrip-thoracy opticine) was prefereded orbenizably and profiled and excitoble determined registerine, state (1989) DRA 3, 401). To 10 00 250 units dissolved in 50 all of water were added 10 all of 1.0 M columbrotis, pir 30, and 500 all of dissillated disemblyformatine containing 20 mg of p-phenytripud distribucyants. The excitor was vorticated and so the 2 for all room temperature in the dark. Approximately 3 ml of houtback was them added. Aller excitoring, adding 3 of all order, and vorticing again, the total was certificated and the sylferide hope theyer described. The estimation process was repeated with subsequent was aller to the contract of the contract of the contraction process was repeated with subsequent execution. The notice of HIPP in 200 all of 1.0 M florates, In 62, was added. The relinder was vorticed

then set at room temperature overnight in the dark.

Separation of the HRP-DNA consignate from their exception and DNA was achieved on a 7% polycarylarised goal the 250 all receions makes was quarted with 100 all of 25% pipced, 15% SSS, 0.5% tomophenol bias, 2.5 mill EDTA. The solution was their distributed into 10 lanes of a 20 x 20 1.15 on mpl and ma at 80 m/hmps under standard conditions (Marchan, A., and Clibert, W. (RIOS) Methods in England SS, 459-550) until the between places of 250 down the gall. The galls were set on Batter FSS4 fills on 50 pitch to that below convoiced with Steam Write (prob) and examined with a batterial to the fill of the set of 150 pitch and 150 p

sodium phosphale, plrt 75, was added, then set at noon temperature overnight.

The contents of the column were filtered through the first at the column bottom into an Amicon Centricon microconcentrator that had been washed twice with distilled water. The HRP-DNA conjugate was then concentrated by contribuyation at \$500 rpm and washed twice with 1x PBS also by centrifugation. The final 1s abulion was then stored at 4°C.

Example 18

Assay for HBV DNA Using HRP-DNA Probe and a Biotinylated Probe Bound to an Avidin Bead

Both labeled proce (6°, 1000 percles in 607.11 of wheth was combined with 5 ml of a 102 % level, solution of 0.8 at a drind hosted filterate bloomateles), 1 ml of 10.8 500, 5.0 ml of 1.5 M/N 60 ml 0.8 ml of 1 mg/ml polys. After 1 h at 37°, 5 the beads were washed before by contribugation with 4c 850, 1.5% 1640 host total or 1.25 ml of the solution. The EV analysis (solvend donew) in 3 sil were was diseased into 1 size of 1.5 ml of

The beefs were watched brice by centrifugation with 4c SSC, 0.1% NP4Q, then taken up is 50 ut of 1% NP4Q, in printing byte, 10 mg/mls SAL XF PSS containing 1 stronds of RRPO-QN conjuspes and set is 37°C for 1 h. The beads were wented with 0.1% NP4Q, XF PSS three times then tendersed in S0 ut to a microtifer cith. To each well, 50 ut it files NPD outdoor files mg OPP (0.2-bring-incidentive), 20 ut 10 to 10 microtifer cith. To each well, 50 ut if then OPD outdoor files mg OPP (0.2-bring-incidentive), 20 ut 10 to 10 microtifer cith. To each well, 50 ut 10 microtifer cith. The description of the outgoing outgoing

Table 4

Condition	Absorbance Reading
1 pmole	>2
0.1 pmole	>2
0.01 pmole	0.88 ± 0.23
1 fmole	0.20 ± 0.05
0.1 fmole	0.07 ± 0.03
NO ANALYTE	0.01 ± 0.01

Cleims

- An assay method for detecting a nucleic acid sequence in a sample, employing two sets of reagents: a first labeling set; and a second capturing set, said method comprising:
 - combining in a liquid medium under binding conditions for complementary pairs, said sample containing analyte in single-stranded form, members of said labeling set of reagents comprising:

- (a) a parality of labeling nucleic acid probes, different probes having a different first analyte complementary sequence and a first noncomplementary region comprising a label reagent recognition sequence, said first analyte complementary sequence having a nucleic acid sequence about 15 to 100 nucleotides in length and said first noncomplementary region optionally being at most about 5 to in length; and
- (b) a labelling reagent having a nucleic acid sequence complementary to said label reagent recognition sequence, wherein said label provides, directly or indirectly, a detectable signal; and members of said second capturing set of reagents comprising.
- (c) a plurality of capturing nucleic acid probes, different probes having a different second analyte complementary sequence and a second noncomplementary region comprising a capturing reagant recognition sequences, said second analyte complementary sequence having a nucleic acid sequence about 15 to about 100 nucleotides in length and said second noncomplementary region optimize behind at most shock 5to in lenetize and.
- (d) a capturing reagent having a nucleic acid sequence complementary to said capturing reagent recognition sequence and capable of binding, directly or indirectly, with a separation means;
 - separating said label into a bound phase and an unbound phase by means of said separation means; and
 - detecting the amount of bound or unbound label as determinative of the presence of said analyte.
- 2. A method scoording to claim 1, wherein eaid nucleic acid sequence complementary to said capturing reagent recognition sequence is bound directly to said separation means, and said analyst., (a), (b), and (c) are combined together for a time sufficient for nucleic acid complement for time formed by the addition of said nucleic acid sequence complementary to said capturing reagent recognition sequence bound to said securation means.
- A method according to claim 1, wherein said separation means is a solid support and wherein said capturing reagent is bound thereto.
- as 4. An assay method for detecting a nucleic acid sequence in a sample, employing two sets of reagents: a first labeling set; and a second capturing set, said method comprising:
 - combining in a liquid medium under binding conditions for complementary nucleic acid sequences, said sample containing analyte in single-stranded form, members of said labeling set of reagents comprising:
 - 6 (a) a plurality of labeling nucleic acid probes including a fluorescer or exceyme label, different probes having a different first analysis complementary segion comprising a label reagent recognition sequence, said first analysis compense having a nucleic acid sequence about 15 to 100 nucleotides in length and said first noncomplementary region optionally being at most about 55 to 100 nucleotides in length and said first noncomplementary region optionally being at most about 55 to 100 nucleotides.
 - (b) a labeling reagant having a nucleic acid sequence complementary to said label reagent recognition sequence; and
 - members of said second capturing set of reagents comprising:
 - (c) a planality of capturing nucleic axid probes, different probes having a different socond enably-complomentary sequence and a second necomplementary region comprising a capturing resport recognition sequence, said second analyte complementary sequence having a nucleic axid sequence about 15 to about 100 nucleotides in length and said noncomplementary region optionally being at most about 50 to about 100 nucleotides in length and said noncomplementary region optionally being at most about 55 to length; and
 - (d) a capturing reagent having a nucleic acid sequence complementary to said capturing reagent recognition sequence;
 - binding said capturing reagant to a solid support through a specific binding pair comprising a hapten and an antibody, either during or after said combining;
 - separating said label into a bound phase and an unbound phase by means of said solid support;
 - detecting the amount of bound or unbound label as determinative of the presence of said analyte.
 - A kit for detecting a nucleic acid analyte comprising:
 (1) members of a labeling set of reagents comprising:

FP 0 225 807 R1

- (a) a plurality of labeling nucleic acid probes, different probes having a different first analyte complementary sequence and a first nucleocomplementary region comprising a label reagent recognition sequence, said first analyte complementary sequence having a nucleic acid sequence about 15 to 100 nucleotides in length and said first noncomplementary region optionally being at most about 5 to it neight; and
- (b) a labeling resgent having a nucleic scid sequence complementary to said label reagent recognition sequence, wherein said label provides, directly or indirectly, a detectable signal; and (2) members of a second capturing set of respents comprising:
- (c) a plarality of capturing nucleic acid probes, different probes having a different second analyse complementary sequence and e second nancomplementary region comprising a second captuing reagent recognition acqueroe having a nucleic acid sequence about 15 to ebut 100 nucleotides in length and said second noncomplementary region optionally being at most about 5 bit in length;
- (d) a capturing reagent having e nucleic acid sequence complementary to said capturing reagent recognition sequence a first member of a specific binding pair conjugate; and
- (a) support meens conjugated to a second complementary member of said specific binding pair.

The use of the following in the manufacture of a kit for detecting a nucleic acid sequence: (1) members of a labeling set of reagents comprising:

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- (e) a plurellity of labelling nucleic ecid probes, different probes having a different first analyse complementary sequence and a first noncomplementary region comprising a label respert recognition sequence, self lett analyse complementary sequence having a nucleic ecid sequence about 15 to 100 nucleotides in length and said first noncomplementary region optionally being at most about 5 to in length; and
- (b) e lobeling reagent having a nucleic acid sequence complementary to said label reagent recognition sequence, wherein said label provides, directly or indirectly, e detectable signet; end
 (2) members of e second capturing set of reagents comprising:
- (c) a plurality of capturing nucleic acid probes, different probes having a different second enables complementary sequence and a second encomplementary region comprising a second cepture of the complete compression of the compression of the
 - оки in erigin; (d) e capturing reagent heving a nucleic acid sequence complementary to said second recognition sequence and containing a first member of e specific binding peir conjugate; end
 - (e) support means conjugated to a second complementary member of said specific binding pair.
 - The use of a polynucleotide probe in an assey method according to any one of claims 1 to 4, wherein the probe has at least two nucleotides, at least one of which is given by the structure

wherein R¹ is a near/we group derivatized with a descrable label, R² is an optional linking moiety including an amide, thiceleter or disulfied integer or a combination thereor, R² is selected from the group consisting of hydrogen, methyl, bromine, fluorine and lodine, R² is H, OH or OR where R is a protecting group and x is an integer in the nance of 1 to 8 fluorine.

8. The use according to claim 6 wherein a polynucleotide probe as defined in claim 7 is employed.

Patentansprüche

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- Testverfehren zum Nachweis einer Nucleinsäuresequenz in einer Probe, wobei zwei Reagenziensätze eingesetzt werden: ein orster Markierungssatz und ein zweiter Einfangsatz, wobei das Verfahren
- folgende Schrifte unflast: Kambinstein in einem Füllseigmedium unter Bindungsbedingungen für komplementlire Paare der Probe, die den Analyt in Einzelstrangform enthält, Bestandleiken des Reagenzienmankerungssatzes, umfassand:
 - (a) eine Veitzähl von Maritierungstrucknisturesonden, unterschiedliche Sonder mit einer unterschiedlichen ersten Analytiensplomentällungenez und einem ersten Nichtlömptiementältbreicht, umtassend eine Marifierungstreugenz-Erkennungssoquenz, wobei die erste Analytiensplomentältsequenzeine Niuciensältursequenz von einer 15 bis 100 Niuchedden Länge hat und der erste Nichtiensplementältbreicht gegebenntstellt bichstrates eine Länge von eine 16 bis dewindt, und
- b) ein Markierungsreagenz mit einer Nucleinsäuresequenz, die komplementär zu der Markierungsreagenz-Erkennungssequenz ist, wobei die Markierung direkt oder indirekt ein nachweisbares Signal liefert: und

Bestandteilen des zweiten Reagenzieneinfangsatzes, umfassend:

(c) eine Vietzell von Enfangruckeinslauresonden, unterschiedlichen Sonden mit einer unterschiedlichen zweiten Analytikomplementilissperaze und einem zweiten Analytikomplementilissperaze und einem zweiten Analytikomplementilissperaze, unterschiedlichen Einfangreagenz-Ertennungssequenz, wobei die zweite Analytikomplementilissperagenz eine Nucleinslauresoquenz von einer 15 bis einer 100 Nucleotiden Länge hat und der zweite Nichtisomplementilissperach opspehendfalls Abbelange eine Längen von einer 5 bis dewiest, und

- (d) ein Einfangreagenz mit einer Nucleinsäuresequenz, die komplementär zu der Einfangreagenz-Erkennungssequenz ist, und das zur direkten oder indirekten Bindung mit einem Trennmittel f\(\frac{1}{2}\)hig ich in der i
- Auftrennen des Markers in eine gebundene und eine nichtigebundene Phase mit Hille des Trennmittels; und Nachweis der Menge an gebundenem oder nicht-gebundenem Marker zur Bestimmung der Gegenwart des Analyten.
- Verfahren nach Anspruch 1, wobei die Nuchiensburesaquenz, die komplementilir zu der Einfangnasgenz-Erkonnungssequenz ist direkt an des Trenomittels gebunden ist, und der Analyt, (a), (b) und (c) für einen Zeitrum zusemmenggeben werden, der zur Bildung der Nuchiensbursendens ausreicht, gefolgt von der Zugabe der Nuckiensbursenquenz, die komplementilir zu der Eintangnagenz-Ersennunssesourseit zitt die an der Tunnmittel debunden.
- Verfahren nach Anspruch 1, wobei das Tronnmittel ein fester Träger ist und wobei das Einfangreagenz deren gebunden ist.
- Testverfahren zum Nachweis einer Nucleinsäuresequenz in einer Probe, wobei zwei Reagenziensätze eingesetzt worden: ein erster Markierungssatz und ein zweiter Einfangsatz, wobei das Verfahren folgende Schritte umfatst.
- o Kombination in einem Flüssigmedium unter Bindungsbedingungen für komplementäre Nucleinsäuressquenzen der Probe, die den Analyt in Einzelstrangform enthält, Bestandseiten des Resgenzienmarkierungssatzen, umfassend:
 - (e) eine Vieizahl von Mantienungsrucieireileaureonden, einschließlich einer Plusreszenz- oder Enzymanitierung, unterschiedliche Sonden mit einer unterschiedlichen orsten Analytiomplementilizereilen, unfassend eine Meritierungsresgenz-Ernernungszegenz, nobei die entst Analytiemplementilizerequinz eine Nucleinsäuresgeparz von dews 15 ib. On Nucleinsäuresgeparzesgepa
- Länge von etwa 5 kb zufweist, und (b) ein Metkenngsresgeza mit einer Nucleinsäuressquenz, die komptementär zu der Merkierungsresgenz-Erkennungssequenz ist und
 - Tougeta Lindmangeologium sie van der Schaffel van der Schaffel van der Schaffel van der Schaffel van Einfangenzieneinfangeatzes, umfassend:

 (c) eine Vielzahl von Einfangenzieneinfangeatzes, umfassend:

 (d) eine Vielzahl von Einfangenzieneinstage von der verschiedliche Sonden mit einer unterschiedlichen zweiten Nichtformielmentlische von der Verschiedlichen zweiten Nichtformielmentlische von der Verschiedliche von der Verschiedliche
- aend eine Einfengresgenz-Ektennungsseguenz, wobel die zwelte Anelyticmplementlinseguenz eine Nuclehstürzeseguenz von einer 15 bis ehre 100 Nucleotiden Linge hat und der Nichtsomplementlinbereich gegebenenfalls Nichtstens eine Linge von etwe 5 kb aufweits, und (d) ein Einfangresgenz mit einer Nucleinstürzessgeunz, die komplementlit zu der Einfangresgenz-
- Erkennungssequenz ist;
 Bindung des Einfangrespenzes an einen festen Träper über ein spezifisches Bindungspar, umfassend
- ein Hagten und einen Antikörper, entweder während oder nach der Kombination; Auftrennen des Markers in eine gebundene und eine nichtgebundene Phase mit Hille des festen Träcers und
- Nachwels der Menge an gebundenem oder nichtgebundenem Marker zur Bestimmung der Gegenwart des Anslyten.
- 5. Kit zum Nachweis eines Nucleinsäureanalyten, umfassend:

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- Bestandteile eines Reagenzienmarkierungssatzes, umfassend;
- (a) eine Vielschil von Kankinnungsmockeischreuworden, unterscheidliche Sorden mit einer unterscheidliche enter anzeite Anzeitenbemenstlänsgenen und einem ertem Knichtinnoprimentlänseringen untersend eine Mehrleungsmegen-Einermagnengartz, wohl die erüb Ausgischreinertiktungsmegen-Einermagnengartz, weber die erüb Ausgischreinertiktungsmegen-Einermagnengartz, eine ein Einge von eine Sie durwiete, und (b) ein Medichrungsmagner mit einer Auschlensührengarte, die komplementik zu der Mesidengungsmegen-Einermagnengenen ist, webei die Mehrleung gelend der überte ein nachmeisten der Versichte ein nachmeisten der Versichte ein nachmeisten der Versichte ein nachmeisten der Versichten der versicht ein nachmeisten der Versichten der Versichten der versichten der versichten der versichten der versichten gelend der versicht ein nachmeisten der versichte den versichten der versichten der versichte den nachmeisten der versichten der versichte ein nachmeisten der versichten der vers
- (2) Bestandteilen eines zweiten Resgenzieneinfangsstzes, umfassend:
 - (c) eine Vielzehl von Einfangnucieinsäuresonden, unterschiedliche Sonden mit einer unterschiedlichen zweiten Analytkomplementärssouenz und einem zweiten Nichtkomplementärbereich, umfagten.

FP 0 225 807 R1

- send eine zweite Einfangreagenz-Erkennungssequenz mit einer Nucleinsäuresequenz von etwa 15 bis etwa 100 Nucleodiden Lillinge, und wobei der zweite Nichtkomptementärbereich gegebenenfalls höckstens eine Länne von etwa 5 ich aufweist.
- (d) ein Einfangreagenz mit einer Nucleinsäuresequenz, die komplementär zu der Einfangreagenz-Erkonnungssequenz ist und einen ersten Bestandheil eines spezitischen Bindungspaarkonjugsts enthält, und
- (e) ein Trägermittel, das an einen zweiten komplementären Bestandteil des spezifischen Bindungspaares konjugiert ist.
- 10 6. Verwendung der folgenden Bestandteile für die Herstellung eines Kits zum Nachweis einer Nucleinsäu-
 - (1) Bestandteile eines Reagenzienmarkierungssatzes, umfassend:
 - (a) eine Voltzahl von Markerungsruudeinsätzereonden, unterschiedliche Sonden mit einer unterschiedlichen serten Ansigktronispensetifiziereiner, der diesen setzen Mohltchreighemerifiziereinet, umfassend eine Markerungsreagenz-Erkensungssoquenz, wobel die erste Analyticemplementlizsequenz eine Modelmatisversequenz von der 15 bis 100 Mohltchodiden Länge het und der orste Nottionsplementlizhereich gegebenerfalls biChistense eine Länge von einen 5 bis aufweit, und
 - (b) ein Markierungsreagenz mit einer Nuclainsäurosoquenz, die kompiementär zu der Markierungsreagenz-Erfannungssaquenz ist, wobei die Markierung direkt oder Indirekt ein nechweisberes Signal liefert, und
 - (2) Bestandteile eines zweiten Reagenzienelnfangsatzes, umfassend:
 - (c) eine Vielzaht von Einfangruschiensturssonden, unterschiedliche Sonden mit eine unterschiedlich chen zweiten Anstytischen gefreigenen dieseugenz und einem zweiten Richtischingenendlichersich, unfassond eine zweite Einfangrasgenz-Eriennungssegunz mit einer Nozleinstursesgunzer von etwe 15 bis etwe 100 Nucleicklien Ellinge, und wobei der zweite Nichtischingenmensführerich gegebe-
- nerfells höchstens eine Länge von eitve 5 kb aufweist.
 (d) ein Eintangreagnach mit einer Nubelmisiturseopustez, die komplementilir zu der zweiten Ertonnungssequenz ist und einen ersten Bestandtell eines spezifischen Bindungspearkonjugets enthält,
- (e) ein Trägermittel, das an einen zweiten komplementären Bestandteit des spezifischen Bindungsperes kontuniert ist.
- Verwendung einer Polynucleotidsonde in einem Testverfahren nach einem der Ansprüche 1 bis 4, wobei die Sonde mindestens zwei Nacleotide aufweist, wobei mindestens eines die folgende Struktur sufweist.

- In der RI eins reaktive Gruppe ist, die mit einem nachweisbaren Marker derivstleiset ist, RI eine naufstalve Verbründungseinheit ist, umfassand eine Amid-, Thioseher- oder Disutifiderinführung oder eins Kombination davon, RP ausgewählt ist aus Wasserstoff, einer Methylinguppe, einem Brom-, Fluor- und loddom, RP ein Wassersfottelom, also OH-Gruppa oder ein OH-Rest ist, wobel R eine Schutzpuppe ist und x eine oausz Zahl im Beracht von einschließich 1 bis diest.
- Varwendung nach Anspruch 6, wobei eine Polynucleotidsonde gem

 ß der Definition in Anspruch 7 eingesetzt wird.

as Revendications

- Procédé d'assai pour détecter una séquence d'acida nucléique dans un échantillon utilisant deux ensembles de réactifs:
- un premier ensemble de marquage : ot un second ensemble de capture, lodit procédé comprenant : la combination dans un militie liquide, dans des condidions de faistion de paires complémentes, dutit échantillon contenant la substance à analyser sous une forme monocaténaire, las éléments dudit ensemble de réscrité de marquage comprenant par la comprenant de la com
 - (a) pixieirus sondes d'acides nucléiques do marquaga, les sondes différentes synt une première esquence complémentaire de la existence à analyser et une première région non complémentaire compronent une esquence de reconnaissance de récald de marquage différentes; Leidis première esquence complémentaire de la subdance à analyser synt une régionne d'acide nucléique d'environ 15 à 100 mucléidates de longueur et ladis promière région non complémentaire eyant facultés-vomment ne focusur d'on plus environ à 51 à .
- (b) un réactif de marquage eyant une séquence d'acide nucléique complémentaire de ladito séquence de reconnaissance de réactif de marquage, où ledit marquaur foumit directement ou indirectement un signal délectable; et
 - las éléments du second ensemble de réactifs de capture comprenant :
- (c) platieurs sondes d'acides nuclifiques de capture, les condes d'ilferintes ayent une socrode dequence complémentaire de la sobatema à analyser en une soccoré égiture no complémentaire compresent une séquence de reconssissance de résert de capture d'ilferiente, solds soccorde séquence complémentaire de la sebetteme à ambigée yeur une séquence d'acide succipies d'environ 15 à environ 100 nuclédides de losqueur et faulte soccorde réglen non complémentaire stant focultairement loques d'use serviers à les environs à les chies de la complémentaire dans focultairement loques d'use serviers à la complémentaire de la complémentaire de la complémentaire destinations de la complémentaire des la complémentaire de la com

FP 0 225 807 R1

- (d) un résctif de capture ayant une séquence d'acide nucléique complémentaire de ladite séquence de reconnaissance du résctif de capture et capable de se lier, directement ou indirectement, avec un élément de séparation :
- la séparation dudit marqueur en une phase liée et une phase non liée au moyen dudit élément de séparation : et
- le détection de la quantité du marqueur lié ou non lié comme détermination de la présence de ladite substance è analyses.
- 2. Procédé seton la revendication 1, dans loqual lastite réquirect d'acide mudélique complémentaire de lastité séquence de racomissaison de résett de captaire est lié directionnet autil étiliser de séparation et le substance à analysex, (a), (b) et (c) cont combinés ensemble prediat un temps suffisent pour qu'îl le forme des compliseuré d'acide mudélique, partie qu'il on ajouts latice séquence d'acide mudélique complémentaire de ladite séquence de recommissance de résettif de capture lide autil étilement de sécuration.
- Procédé selon le revendication 1, dans lequal tedit élément de séparation est un support solide et ledit récotif de capture lui est lié.
- Prooféé d'assai pour détecter una séquence d'acide nucléique dans un échanition utilisant deux ansambles de réactifs:
- un premier ensemble de marquaga; et un second ansemble de capture, ledit procédé comprenant : le combinaison, dans un milleu liquide dans des conditions de liaison de séquencee complémental
 - res d'ecides nucléliques, dudit échantillon contanant la substance à analyser sous une forme monocaténaire, les éléments d'udit ansemble de récrité de marquega compranant :
- 6 (a) plasiours sondes d'acides nucléiques de marquage comprenent un marques fluorecant ou anymetique, les condes différentes eyent une premise eliquance complimentaire de la reconstance à embyers et une premitre eliquine complémentaire comprenent une séquence de reconsissance de de récell de marquage différente ; lestes premitre eliquines complémentaire de les techniques et de la marque d'inférente ; lestes premitre eliquines complémentaire de les techniques et eliquines et perfect de la marque d'inférente ; lestes premitres despuerce ou confidence de la marque d'inférente en les forces d'un ben enforte 5 foi uniférent en la forces d'un ben enforte 5 foi uniférent elle promité et de la marque d'un ben enforte 5 foi uniférent elle promité d'un ben enforte 5 foi uniférent elle promité de la marque d'un ben enforte 5 foi uniférent elle promité de la marque d'un premitre de la marque de la
- pramièra région non complémentaire eyent facultétévement une longueur d'au plue environ 5 kb; et (b) un réactif de marquise avent une efiquence d'acide nudifique complémentaire de la dite séquence de reconnaissance de réactif de marquage;
 - las élémants du second ensemble da réactifs de capture comprenant :
- (c) justissur sondas d'acides nuclifiques de captur, les sondes différentes syaru une seconde séquence concilientative de la bestance à ansigne et une seconda région non complémentaire comprenant une séquence de racomaissance de résidif de capture différents, lacife seconde séquence complémentaire de la substance à surjever syert une séquence d'acide nuclidique d'arviron 15 à envienn 100 nuclésidas de longueur at tades région non complémentaire étant focultais-mante floque d'as plus anviron 16 à re.
- (d) un réactif de capture ayant une séquence d'acide nucléique complémentaire de ladite séquence de reconneissance du réactif de capture;
 - la liaison dudit réactif de capture à un support sollide vita une paire à liaison spécifique comprenant un hapithe et un antisonps soit pendant, soit après ladite combinaison; ; la séparation dudit marqueur en une chase liée et une plases on liée au moven dudit succort
 - solide; et la détaction de la quantité du marqueur lié ou non lié comme détermination de la présence da ladits substance à analyser.
- so 5. Nécassaira pour la détection d'una substance à analyser constituée d'acides nucléiques comprenant :
 - (1) les éléments d'un ensembla de réactifs de marquage compresant : de plusiours sondes d'acides nucléiques de marquage, les sondes différentes ayant une première séquence complémentaire de la substance à analyser et une première région non complémentai-
- re comprenant une sidquence de reconnaissance de récutif de marquage différentes; i subte première séquence complémentaire de la substance à analysera system use séquence d'acides nucléiques d'environ 15 à 100 nucléoides de longueur et lacite première région non complémentaire avant faculité-ownent une lonnauer d'au alus aveniron 5 lés : et

EP 0 225 907 R1

- (b) un réactif de marquage ayant une séquence d'acide nuclétique complémentaire de la dite séquence de réactif de marquage, où ledit marqueur fournit directement ou indirectement un sinnal référetable : et
- (2) les éléments d'un second ensemble de réactifs de capture comprenant :

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- (c) plusivers conduc d'exident musifiques de capture, les condes différentes ayant une seconde réglament complimitation de la settation à analyses et ne seconde réglam non complémentaire me, compresent une seconde séquence de reconneissance de résield de capture, dufférentes, la complexión de la capture, de la capture de la substracta à analyses ayant une séquence d'action la capture à analyses de la capture de la substracta à analyses ayant une séquence d'action una capture d'entre et la capture de la substracta à analyses ayant une séquence d'action complémentaire sinte facultativement la capture des des parties de la capture de l
- (d) un réactif de capture ayant une séquence d'acide mucléique complémentaire de ladite séquence de reconsissance du réactif de capture et un premier élément d'un conjugué de paire à liaison apécifique; et
- (e) un élément support conjugué à un socond élément complémentaire de ladite paire à liaison spécifique.
- Utilisation des composants sulvants dans la fabrication d'un nécessaire pour détecter une séquence d'acide nucléique :
- (1) les éléments d'un encemble de récolts de marquage compresant : (a) plusieurs sondes d'accides nucléiques de marquage, les sondes différentes ayant une première séquence complémentaire de la substance à sentyers et une première région non complémentaire re compresant une séquence de reconnaissance de récolf de marquage différentes ; lacite première séquence complémentaire de la substance à écalité en sant pas de la complémentaire de la substance à la califer evant une séquence d'actide
 - nuclétique d'environ 15 à 100 auclécitates de longueur et tadite première région non complémentaire syent facultativement une longueur d'au plus environ 5 ls ; ot (b) un réactif de marquage ayant une séquence d'acide nuclétique complémentaire de le dite
- séquence de reconnaissance de réactif de merquage, où ledit marqueur fournit directement ou indirectement un aignat détectable ; et (2) les éléments d'un second ensemble de réactifs de capture comprenant :
- (c) platificars acostos d'acidose nuclátiques de capture, les condes différentes ayent une seconde séquence complémentaires de la substance à avaigner et une seconde région non comprémentaine comprenent une seconde séquence de recomaissance de récett de cepture, différentes, lettre seconde séquence complémentaire de la sobratione à analyser eyet une séquence disnuclifique d'environ 15 à anviron 100 nucléditées de langueur et ladite seconde région non complémentaire deatré trudutement intruse d'aux servierres (54 or 1).
 - (d) un réactif de capture ayant une séquence d'acide nucléfique complémentaire de ladite seconde séquence de réconsaissance et contenant un premier élément d'un conjugué d'une pair à laison solécitique : et
- (e) un diément support conjugué à un second élément complémentaire de ledite paire à liaison spécifique.
- Utilisation d'une sonde polynuciéotidique dans un procédé d'essai selon l'une quelconque des revendications 1 à 4, où la sonde a au moins deux nucléotides dont au moins un répond à la structure

dans lequelle R² est un groupe réactif dérivétée avec un marqueur défectable, R² est un fragment leur feculteit comprenent une liaison améle, théofère un distinute, ou une de leurs combinésone, R² est chôid dans le groupe constitué par un aisme d'hydrogène, de brence, de fibre ou d'inde us groupe méthyle, R² est H, OH ou OR, où R est un groupe protecteur et x est un enfer dans la gemme de 1 à 8 inclusivement.

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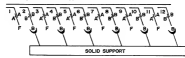
 Utilisetion selon la revendication 6, dans laquelle on utilise une sonde polynuciéctidique telle que définie dans la revendication 7.

COMPLEXES FORMED WITH ANALYTE

i) Using DNA solid support



2) Using Avidin solid support



→ = Avidin

IA-3A-5A-7A-9A-IIA= Labelling set 2B-4B-6B-8B-IOB-i2B= Capturing set FIG. I